PATENT SPECIFICATION

613,782



Convention Date (United States of America): July 11, 1945.

Application Date (in United Kingdom): July 2, 1946. No. 19808 46.

Complete Specification Accepted: Dec. 2, 1948.

Index at acceptance:—Class 2(iii), D1t.

COMPLETE SPECIFICATION

Phthalocyanine Sulfonhydrazides

We, General Aniline & Film Cor-PORATION, a corporation organized and existing under the laws of the State of Delaware, United States of America, of 5 230, Park Avenue, New York, County and State of New York, United States of America (Assignees of ARTHUR LAWRENCE Fox, a citizen of the United States of America, of 118, Parker Avenue, Easton, 10 County of Northampton, Commonwealth of Pennsylvania, United States of America), do hereby declare the nature of this invention and in what manner the same is to be performed, to be particu-15 larly described and ascertained in and by the following statement:-

This invention relates to new phthalo-

cyanine derivatives.

The term phthalocyanine is used 20 herein in the generic sense to define tetraazaporphins in which each of the four pyrrole nuclei is fused to an aromatic nucleus, e.g., phenyl, diphenyl, naphthyl, anthranyl, etc., of which phthalocyanine itself (tetrabenzotetraazaporphin) is a well-known example. The phthalocyanine molecule may be metal free or contain a metal in complex combination, for 30 example, copper, cobalt, nickel, iron and aluminium.

The invention concerns new compounds \mathbf{which} which are phthalocyanine sulfon-hydrazides and the preparation thereof mydrazides and the preparation thereof
which may be accomplished by condensing a phthalocyanine sulfonylchloride with a hydrazine which may be
hydrazine itself or a primary or
secondary alkyl, substituted alkyl,
alicyclic, aryl or heterocyclic hydrazine.
All of these hydrazines contain a
reactive hydrazen atom attached to one reactive hydrogen atom attached to one of the nitrogens of the -N.N- group.

The preparation of the new phthalo-45 cyanine derivatives can be effected by reacting the corresponding phthalocyanine sulfonylchloride with the hydrazine at room or elevated temperatures in the presence of a suitable

reaction diluent, for example, water, 50 alcohol, acetone, nitrobenzene or an excess of the hydrazine.

In addition to hydrazine itself, suitable hydrazines for the preparation of the new compounds are, for example, 55 methyl, ethyl, propyl and butyl hydrazine, sym. and asym. dimethyl hydrazine, N,N - bis(β - hydroxyethyl) hydrazine, cyclohexyl hydrazine, benzyl hydrazine, cyclonexyl nydrazine, benzyl hydrazine, phenyl hydrazine, p-chloro-60 phenyl hydrazine, p - bromophenyl hydrazine, p-sulfophenyl hydrazine, p-sulfophenyl hydrazine, p-xylyl hydrazine, 3-hydrazinoaniline, sym. and asym. diphenyl hydrazine, triphenyl 65 hydrazine, naphthyl hydrazine, a-methylphenyl hydrazine, a ethyl phenyl hydrazine, a, dimethyl phenyl hydrazine, a, bydrazine, a-lepidyl hydrazine, 70 a-hydrazoquinoline, 4-aminomorpholine, 3-hydrazino - 1,2,4-triazole and 5hydrazinotetrazole.

The new products are phthalocyanines containing attached to the arylene 75 nuclei one or more unsubstituted or alkyl, substituted alkyl, alicyclic, aryl or heterocyclic sulphonhydrazide groups. In addition to the sulfonhydrazide group or groups, other substituents, such as 80 halogen atoms, e.g., chlorine, or amino or carboxy radicals may be attached to the arylene nuclei of the phthalocyanines. The number of the sulfon-hydrazide groups on the arylene nuclei 85 may be varied depending on the number of sulfonylchloride groups on the arylene nuclei of the corresponding phthalo-cyanine sulfonylchlorides. In some instances the new products may also 90 contain attached to the arylene nuclei one or more sulfonic acid groups, either free or as the hydrazine salt, not all of the sulfonylchloride groups being converted to sulfonhydrazide groups in the 95 reaction. They range in color from bluish-green to green, being somewhat greener in shade than the phthalo-

cyanine sulfonamides described in Specification No. 520,199. They may be employed as pigments or as dyestuffs depending on their solubility

characteristics.

Dyestuffs which are soluble in dilute aqueous caustic alkali, e.g., sodium, potassium or lithium hydroxide, may be prepared by introducing onto the arylene 10 nuclei of the phthalocyanine molecule a plurality of sulfonhydrazide groups containing a free hydrogen atom on the nitrogen of the sulfamyl portion thereof. Compounds of this character are capable 15 of dyeing directly cotton, wool, and silk, showing particularly good affinity for silk. Employed in the form of their particularly salts, soluble alkalinepotassium, ammonium sodium,

20 lithium, they are suitable as colorants for the preparation of writing ink fluids, showing to particular advantage in writing inks rendered quick drying by small amounts of caustic alkali, being, 25 generally, quite stable in dilute aqueous caustic alkali.

The starting phthalocyanine sulfonyl-chlorides employed in the preparation of the phthalocyanine sulfonhydrazides may 30 be obtained by reacting chlorosulfonic acid with the corresponding phthalocyanines at an elevated temperature as described in Specification No. 515,637. They may also be made from the corre-35 sponding phthalocyanine sulfonic acids or their salts by treatment with

phosphorus pentachloride. The invention is further illustrated by

the following specific examples to which, 40 however it is not to be limited. Parts are by weight unless otherwise noted.

EXAMPLE 1.

Ten parts of hydrazine hydrate is dissolved in 50 parts of water and 10 parts 45 of copper phthalocyanine tetrasulfonylchloride added. Stirring is then instituted and continued for several hours at room temperature until dis-The reaction solution is complete. 50 mixture is then stirred into 4 volumes of 10% hydrochloric acid and the precipitated dyestuff filtered and dried. There is obtained a bluish-green powder which is quite soluble in dilute aqueous sodium 55 hydroxide. The yield is 89.3%.

EXAMPLE 2.

Twenty parts of copper phthalocyanine tetrasulfonylchloride is stirred into an excess of phenyl hydrazine which is 60 heated on the steam bath. The mixture at once begins to foam and the reaction takes place quite rapidly. In 30-45

minutes only a dark green solid material remains. This material is then stirred into approximately 4 volumes of 10% hydro- 65 chloric acid and the precipitate filtered and dried. The product thus obtained is somewhat soluble in dilute aqueous sodium hydroxide and dyes silk a vivid green shade. The yield is 90.4%.

Example 3.

Twenty parts of copper phthalocyanine tetrasulfonylchloride is stirred into a mixture of 30 parts of 4-sulfophenyl-hydrazine in 100 parts of nitrobenzene. 75 The resulting mixture is heated to 50— 60° C. while stirring and then allowed to slowly cool to room temperature after which stirring is continued for several hours. The nitrobenzene is then removed 80 by means of steam distillation and the resulting aqueous suspension acidified with 10% hydrochloric acid. The precipitated product is filtered and dried. A green powder is obtained which is quite 85 soluble in dilute aqueous sodium hydroxide and is quite satisfactory for use as a dyestuff for writing ink fluids. The yield is 72%.

Having now particularly described and 90 ascertained the nature of our said invention and in what manner the same is to be performed, we declare that what we

claim is:-

1. A process of preparing a phthalo- 95 cyanine sulfonhydrazide which comprises condensing a phthalocyanine sulfonylchloride with a hydrazine.

2. The process as defined in claim 1, wherein the hydrazine is unsubstituted 100

hydrazine.

3. The process as defined in claim 1, wherein the hydrazine is an alkyl hydrazine.

4. The process as defined in claim 1, 105 wherein the hydrazine is an alicyclic

hydrazine.

5. The process as defined in claim 1. wherein the hydrazine is an aryl hydrazine.

6. The process as defined in claim 1, wherein the hydrazine is a phenyl

hydrazine.

7. Phthalocyanine sulphonhydrazides whenever prepared by a process of any of 115 the preceding claims or by a process which is the obvious chemical equivalent thereof.

Dated this 2nd day of July, 1946. CRUIKSHANK & FAIRWEATHER,

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Leamington Spa: Printed for His Majesty's Stationery Office, by the Courier Press .-- 1948. Published at The Patent Office, 25, Southampton Buildings, London, W.C.2, from which copies, price 2s. 0d. each (inland) 2s. 1d. (abroad) may be obtained.